Effects of attached bacteria on organic aggregate settling velocity in seawater

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ABSTRACT: We investigated whether attached bacteria affect the settling velocity of organic aggregates via modifications of the physical properties of aggregates, including density, porosity, and morphology. Model aggregates, prepared by mixing 2 different polysaccharides (fucoidan and chitosan), were incubated in coastal seawater passed through either 0.8 μm (AGG0.8) or 0.2 μm (AGG0.2) filters. After incubation for 48 h, AGG0.8 were much more densely (3.2- to 10.1-fold) colonized by bacteria than AGG0.2. Based on median settling velocities (\(W_{50}\)), as determined by laser in situ scattering and transmissometry, the \(W_{50}\) of AGG0.8 was lower (1.6- to 4.5-fold) than that of AGG0.2 for a size class of 62 to 119 μm. Stokes model analyses indicated that this reduction in \(W_{50}\) could be largely attributed to the higher porosities of AGG0.8 (0.932−0.981) than those of AGG0.2 (0.719−0.929). Our results support the notion that the modification of aggregate structure by attached bacteria (porosity enhancement) can be an important factor controlling the settling velocity of marine particles.

KEY WORDS: Bacteria · Aggregate · Settling velocity · Porosity · Marine environments

INTRODUCTION

Organic aggregates, including transparent exopolymer particles (TEP) and hydrogels, play an important role in the regulation of material cycling and food webs in marine ecosystems (Alldredge 1979, Simon et al. 2002, Verdugo 2012). Organic aggregate settling from the sunlit layer to the deeper layers of the oceans mediates the vertical delivery of organic carbon and other bioelements, representing an important regulatory element of carbon sequestration (Fisher et al. 1991, Passow & De La Rocha 2006, De La Rocha & Passow 2007) and a means of energy and nutrient supply to heterotrophic organisms in aphotic layers and on the seafloor (Alldredge 1979, Grossart et al. 1998, Passow et al. 2001). Because the settling velocity of aggregates is a key variable that influences the length-scale of vertical carbon delivery and remineralization depth (Passow & Carlson 2012), it is important to clarify the variability and control of aggregate settling velocities in marine environments.

Aggregate settling velocity is affected by the physical properties of aggregates including size, porosity, morphology, and association with ballast particles such as diatom frustules and coccolith shells (Ploug et al. 2008, Iversen & Ploug 2010). These aggregate properties may reflect not only the composition of, and physicochemical interactions among, polymeric materials and other constituents (Chin et al. 1998, Verdugo 2012) but also the modification of aggregate structures by attached microbes. Bacteria can actively proliferate on the surface and in the interstitial spaces of aggregate matrices, cleaving polymer chains to create pores and reduce aggregate size (Passow & Alldredge 1994, Azam & Malfatti 2007). Alternatively, bacteria may excrete sticky polymeric materials to fill interstitial spaces and promote aggregate production (Cowen 1992, Stoderegger & Herndl 1998, Sugimoto et al. 2007). Furthermore, the replacement of aggregate matrices by bacterial cells may alter bulk aggregate density. However, the prevailing mode and net outcome of these complex bac-
terial actions in controlling organic aggregate settling velocities in seawater are not entirely clear.

To understand how bacterial modification of the physical properties of aggregates affects aggregate settling velocities, we examined the settling velocities of model aggregates with different degrees of bacterial colonization. Data on the physical parameters of aggregates, including size, porosity, and morphology, were collected to investigate the dominant mechanism by which attached bacteria affect the settling velocities of organic aggregates in marine environments.

MATERIALS AND METHODS

Preparation of model aggregates

We used compositionally defined, spontaneously assembled gels (Verdugo 2012) as model aggregates to investigate bacterial effects on aggregate settling velocity with minimal effects of compositional variability of organic aggregates in natural environments (see 'Discussion'). The model aggregates were prepared using fucoidan (a sulfated polysaccharide found in brown seaweed) and chitosan (a partially deacetylated derivative of the amino polysaccharide, chitin) according to a gel preparation protocol originally developed by Nakamura et al. (2008) for pharmaceutical purposes. Fucoidan was extracted from *Kjellmaniella crassifolia* (Gagomekombu) according to Nakamura et al. (2008). Gagome-kombu powder (10 g) (Toyonaka Matumae Konbu Honpo Corp., Osaka, www.kobuya.net) was added to 100 ml 70% by volume ratio (v/v) ethanol containing 0.1 M acetic acid in a 200 ml conical glass flask. After gentle mixing, the suspension was filtered through polycarbonate filters (10 μm pore size, 47 mm diameter; Whatman). The filtrate was air-dried and added to 200 ml 8.8 μM CaCl₂ solution. After boiling for 30 min, the extract was centrifuged (2330 × g) at room temperature for 10 min. The supernatant was filtered through polycarbonate filters (10 μm pore size, 47 mm diameter; Whatman) and transferred to a 500 ml conical glass flask. The aggregate suspension was filtered through polycarbonate filters (10 μm pore size, 47 mm diameter; Whatman). The filtrate was mixed with 2x volume of 99.5% ethanol (total volume: ca. 150 ml) and centrifuged (2330 × g) at room temperature for 10 min. Finally, the precipitates (fucoidan extracts) were broken into small pieces with spatulas and air-dried in a flow hood. Fucoidan solution (2.1% wt/vol) was prepared by dissolving 16.6 mg fucoidan extract in 790 μl Milli-Q water (Millipore). Chitosan solution (3.8% w v⁻¹) was prepared by dissolving 1.25 g chitosan (LL-40, Yaizu Suisankagaku Industry, extracted from *Chionoecetes opilio*; nominal deacetylation rate: >80%) in 33.1 ml 190 mM HCl.

Model aggregates were prepared by mixing chitosan and fucoidan solutions according to Nakamura et al. (2008) with the following modifications. Before the preparation of aggregates, chitosan and fucoidan solutions (see above) were centrifuged at 1200 × g for 10 min at 23°C. The supernatant of each solution was successively filtered through 0.8 μm and 0.2 μm syringe filters (25 mm diameter, Acrodisc syringe filter with Supor® membrane; Pall) to remove particles. Then, 210 μl chitosan solution was added to 50 ml filtered (0.2 μm) seawater in a 50 ml polypropylene tube (BD Falcon Conical Centrifuge tube; Fisher Scientific). After gentle mixing, 790 μl clear fucoidan solution was added to the seawater and then mixed. At this stage, the formation of aggregates was observed. After storage of the suspension at 4°C for 3 h, aggregates that settled at the bottom of the tube were collected by removing the supernatant by aspiration (~10 ml solution remaining). Then, 40 ml of the filtered (0.2 μm) seawater was added to the tube. The aggregate suspension was filtered through a nylon mesh (Nitex screen, mesh size 200 μm; Wildlife Supply Company) to remove large aggregates. The aggregate suspension (hereafter, 50 ml working suspension) was stored at 4°C for 3 h until it was used for the settling velocity experiments.

Collection of seawater samples and incubation of model aggregates

Surface seawater samples were collected using a plastic bucket at the shore of Oarai Beach, Ibaraki, Japan (36° 19′ 3″ N, 140° 35′ 29″ E), on January 30 (water temperature: 10.1°C, salinity: 29.2), August 29
(24.0°C, 29.2), and November 15 (17.3°C, 28.8), 2011. Within 12 h of sampling, seawater samples were filtered through 0.8 μm polycarbonate filters (47 mm diameter; Whatman) to eliminate grazers and other eukaryotes. The filtrate was further filtered through 0.2 μm polycarbonate filters (47 mm diameter; Whatman) using negative pressure (<20 mmHg) to reduce the abundance of bacteria and minimize their effects on aggregate properties.

The 50 ml working suspension of aggregates and 500 μl autoclaved 20 mM Na$_2$HPO$_4$ solution were added to three 500 ml polycarbonate bottles (Nalge Nunc International) filled with 450 ml seawater (filtered at 0.8 μm or 0.2 μm). After incubation for 48 h at 23°C in the dark using an incubation shaker (TA-20RLS, Takasaki Scientific Instruments; shaking speed: 100 rpm), both aggregate preparations (those obtained from 0.8 μm and 0.2 μm filtration; hereafter, AGG$_{0.8}$ and AGG$_{0.2}$, respectively) were used for the determination of aggregate settling velocities and the characterization of aggregate properties.

**Microscopic examination of aggregates and attached bacteria**

Aggregate samples were fixed with 0.2 μm filtered 2% v/v (final concentration) glutaraldehyde (20% glutaraldehyde; Wako). In the January experiment, the aggregate size and abundance of attached bacteria were determined after SYBR Gold-Alcian Blue double staining (Sugimoto et al. 2007). Seawater samples were filtered through 0.02 μm aluminosilicate filters (25 mm diameter; Whatman). Filters were stained with 1% wt/vol Alcian Blue (8GX; Sigma-Aldrich) dissolved in 3% v/v acetic acid (Logan et al. 1994) and then stained with SYBR Green I (0.25% v/v, in a ratio of 1:400 of the supplied concentration diluted with Milli-Q filtered at 0.2 μm; Molecular Probes; Noble & Fuhrman 1998) for 10 min. The filters were mounted on slides using immersion oil (nd = 1.516; Olympus) and observed under an epifluorescence microscope (×1000, BX61, Olympus) equipped with epifluorescence optics optimized for SYBR Green I (filter set, U-MNIB2; Olympus). After imaging, the volumes of bacteria (μm$^3$) attached to aggregates were determined for AGG$_{0.8}$ and AGG$_{0.2}$ from both their area and perimeter using ImageJ software (Massana et al. 1997).

In the August and November experiments, the abundance of attached bacteria was determined by the 4’,6-diamidino-2-phenylindole (DAPI) staining method (Porter & Feig 1980). Samples were filtered through 12 μm polycarbonate filters (25 mm diameter, Whatman) with a vacuum of <20 mm Hg. The filters were stained with DAPI (1 μg ml$^{-1}$) for 5 min, mounted on slides using immersion oil (Vectashield, from Vector Laboratories, and from Citifluor Limited), and observed under an epifluorescence microscope (×1000, BX61, Olympus) equipped with epifluorescence optics optimized for DAPI (filter set, U-MWU2, Olympus). The sizes of aggregates, detected as pale-yellow objects, were manually determined using a grid ocular micrometer and classified into the 5 size classes described above.

**Abundance, size, and settling velocity of aggregates**

Particle size distribution and particle volumes were measured by laser in situ scattering and transmissometry (LISST, LISST-100X, Sequoia Scientific; Agrawal & Pottsmith 2000) (hereafter, aggregate ‘size’ refers to the ESD of the aggregate). The settling velocities of aggregates were determined using a settling column (300 ml volume, 15 cm travel path) attached to the chamber of the LISST-100X (Small Volume Horizontal Test Chamber; Sequoia Scientific) (Agrawal & Pottsmith 2000). The aggregate settling velocity was estimated from the decrease in aggregate abundance over time. The aggregate abundance data were collected every 30 s (up to 3 h) for each size class (the numbers of aggregates counted at time zero were 841 to 21 693 for each size class) to determine the time ($T_h$, in seconds) at which the aggregate abundance became half of the initial aggregate abundance for a given size class. Then, the median settling velocity ($W_{50}$, cm s$^{-1}$, Owen 1971) was calculated by dividing the travel path (15 cm, the distance between the surface of sample seawater and the horizontal position of the LISST detector) by $T_h$ (range: 900–8060 s).
Density of attached bacteria

In the August experiment, the density distribution of bacteria attached to aggregates (after 48 h incubation) was measured according to Inoue et al. (2007). To remove attached bacteria from aggregate matrices, aggregate structures were mechanically destroyed by filtration through a 0.8 μm syringe filter (25 mm diameter; Pall). First, 2 ml AGG_{0.8} was filtered through the syringe filter, and the filtrate was discarded (we assumed that bacteria collected on the 0.8 μm filter represented those associated with AGG_{0.8}, although some attached bacteria might have passed through the filters). Then, 2 ml filtered seawater (0.2 μm) was passed through the same syringe filter. The filtrate was passed a second time through the same syringe filter. This operation was repeated 10 times to obtain a bacterial suspension.

The recovery of attached bacteria was calculated as follows: (bacterial abundance determined for the bacterial suspension obtained after aggregate mechanical destruction) / (attached bacterial abundance initially determined using the 0.8 μm pore size filters) × 100. Bacteria were enumerated by the DAPI staining method as described in ‘Microscopic examination of aggregates and attached bacteria’. The recovery was 102.1 ± 16.8% (mean ± standard error [SE], n = 7).

To determine attached bacterial density, 1 ml attached bacterial suspension and a 0.1 ml solution of density marker beads (Amersham Biosciences) were layered on top of 9 ml Percoll gradient working solution. The working solution consisted of 30% v/v Percoll, 10% v/v of 10× phosphate-buffered saline (pH 7.4), 2.34% wt/vol of NaCl and autoclaved Milli-Q (filtered through the 0.2 μm pore size filter) × 100. Bacteria were enumerated by the DAPI staining method as described in ‘Microscopic examination of aggregates and attached bacteria’. The recovery was 102.1 ± 16.8% (mean ± standard error [SE], n = 7).

Aggregate porosity

The porosities of AGG_{0.2} and AGG_{0.8} obtained in the November experiment were estimated by comparing aggregate size distributions determined by LISST to those determined using a resistive pulse particle counter (Coulter counter; Multisizer II, Beckman Coulter) (Sterling et al. 2004). This method is based on different principles of 2 instruments to measure particle size distributions. LISST measures light scattering of particles (reflecting particle bulk volume), whereas a Coulter counter determines the electric resistance of particles (reflecting particle solid volume). For porous particles, the particle size distribution determined by a Coulter counter deviates to smaller size categories relative to that determined by LISST, reflecting the degree of porosity (Sterling et al. 2004). We used the following equation to derive the index of porosity (ε):

\[
\text{Porosity}(\varepsilon) = 1 - \frac{V_C}{V_L}\]

where \(V_C\) and \(V_L\) are the volume of aggregates (ml aggregate\(^{-1}\)) determined by the Coulter counter and LISST, respectively. \(V_C\) and \(V_L\) for each size class were estimated from the Coulter-based aggregate diameter (\(D_C\)) and LISST-based diameter (\(D_L\)), assuming that the aggregates were spheres. Because the bins differed between \(D_C\) and \(D_L\), we derived the abundance-\(D_L\) regression (log-log regression) for the LISST data to estimate the aggregate abundance (l\(^{-1}\) μm\(^{-1}\)) for the size category between 62 μm (\(N_{62}\)) and 119 μm (\(N_{119}\)). Then, \(D_C\) for the abundance range between \(N_{62}\) and \(N_{119}\) was compared to the \(D_L\) (derived from the abundance-\(D_L\) regression) belonging to the same abundance category.

Ectoenzymatic activity

In the August experiment, the activity of the bacterial ectoenzyme that potentially hydrolyzes aggregate polymer constituents (chitosan) was determined using a methylumbelliferyl (MUF) substrate,
4-MUF-N-acetyl-β-D-glucosaminide (Sigma-Aldrich) (Hoppe 1983). Fluorescent substrate dissolved in 2-methoxyethanol was added to either total or syringe-filtered (0.8 μm) sample waters (final conc. 0.05 mM). The samples were incubated at 23°C for 2 min. After adding borate buffer (0.5 M boric acid with NaOH, pH 10), fluorescence (455 nm emission, 365 nm excitation) was determined using a fluorescence spectrophotometer (RF-1500; Shimadzu). The enzyme activity was calibrated using 4-methylumbelliferone (Sigma-Aldrich) as a standard. The enzyme activity associated with each aggregate (AGG0.2 and AGG0.8) was estimated by subtracting the enzyme activity determined for filtered samples from that determined for the total sample.

RESULTS

Aggregate size distribution and total volume

After incubation for 48 h, both AGG0.8 and AGG0.2 displayed a similar unimodal size distribution with a peak at 119 μm in all experiments (Fig. 1). Total volume concentration of aggregates initially added was 97.5 ± 4.3 ppm (mean ± SE, n = 22), which was within the range of typical TEP volume concentrations reported in coastal waters (Passow 2002). Aggregates belonging to the 62 to 119 μm size class were used to assess the effect of bacterial attachment on aggregate settling velocities. This size range was used because for smaller aggregates (<62 μm size class) the settling velocity was too low to be determined within the time frame of our measurement (up to 3 h) and because LISST could not capture sufficient numbers of settling events for larger aggregates (>119 μm size class).

Abundance and density of attached bacteria, and settling velocity

Aggregates were porous (Fig. 2a), clearly stained by Alcian Blue (Fig. 2b), and associated with bacter-
ria (Fig. 2c). For both AGG\textsubscript{0.2} and AGG\textsubscript{0.8}, bacterial abundance per aggregate generally increased with increasing aggregate size (Fig. 3). Empirical equations that relate attached bacterial abundance (ABA) to the diameter of aggregates (D in µm) were obtained using a linear regression model after a double logarithmic transformation. The following regression equations were derived: log ABA = -1.32 + (2.31 \times \log D) (r^2 = 0.98, p < 0.001, n = 15) for AGG\textsubscript{0.8} and log ABA = -1.51 + (2.04 \times \log D) (r^2 = 0.57, p < 0.01, n = 15) for AGG\textsubscript{0.2}. For each size class, bacterial abundance associated with AGG\textsubscript{0.8} was much higher (3.2- to 10.1-fold) than the corresponding value for AGG\textsubscript{0.2}. The differences in bacterial abundance between AGG\textsubscript{0.8} and AGG\textsubscript{0.2} were significant (p < 0.01, Student’s t-test) for all size classes. In the January experiment, the average size of bacteria associated with AGG\textsubscript{0.8} was 0.47 ± 0.01 µm\textsuperscript{3} (mean ± SE, n = 474), which did not differ significantly (p > 0.05, Student’s t-test) from the average size of bacteria associated with AGG\textsubscript{0.2} (0.46 ± 0.02 µm\textsuperscript{3}; n = 181). The weighted average density of attached bacteria, determined for AGG\textsubscript{0.8} collected in the August experiment, was 1.064 ± 0.002 g cm\textsuperscript{-3} (n = 4). Note that this estimate has an error due to the inclusion of bacteria in either top or bottom fractions of the density gradient (these bacteria have densities either lower or higher than the analytical limits; see ‘Materials and methods’).

While the abundance of bacteria contained in the top fraction accounted for only 3.7 ± 1.6% (n = 4) of total attached bacterial abundance, the contribution of bacteria contained in the bottom fraction was relatively large (38.1 ± 7.1% [n = 4] of total attached bacterial abundance), suggesting that our estimate of average bacterial density might be too low.

The ranges of \(W_{50}\) values for AGG\textsubscript{0.8} and AGG\textsubscript{0.2} were 0.0025–0.0054 cm s\textsuperscript{-1} and 0.0056–0.0136 cm s\textsuperscript{-1}, respectively. For each size class, the \(W_{50}\) values for AGG\textsubscript{0.8} were lower (1.6- to 4.5-fold) than those of AGG\textsubscript{0.2} (Fig. 4). The differences in the \(W_{50}\) values between AGG\textsubscript{0.8} and AGG\textsubscript{0.2} were significant (p < 0.05, Student’s t-test), except for the 62, 73, and 86 µm size classes in the August experiment (Fig. 4).

**Aggregate \(\epsilon\), enzyme activity, and morphological parameters**

The \(\epsilon\) of AGG\textsubscript{0.8} and AGG\textsubscript{0.2} varied in the ranges of 0.932–0.981 and 0.719–0.929, respectively. The average \(\epsilon\) of AGG\textsubscript{0.2} was significantly smaller than the corresponding \(\epsilon\) of AGG\textsubscript{0.8} for each size class (p < 0.05, Student’s t-test, Fig. 5). Empirical equations that relate \(\epsilon\) to D were obtained using a quadratic regression model after a double logarithmic transformation. A quadratic rather than linear regression model was used because the \(r^2\) value of the quadratic regression exceeded that of the linear regression for both AGG\textsubscript{0.8} and AGG\textsubscript{0.2}. The following regression equations were derived: log \(\epsilon\) = 1.214 + (0.719 \times \log D) − [0.166 \times (\log D)^2] (r\textsuperscript{2} = 0.73, p < 0.001, n = 20) for AGG\textsubscript{0.8} and log \(\epsilon\) = −2.704 + (4.409 \times \log D) − [1.040 \times (\log D)^2] (r\textsuperscript{2} = 0.76, p < 0.001, n = 20) for AGG\textsubscript{0.2}.

In the August experiment, the ectoenzymatic activity (hydrolysis rate of 4-MUF-N-acetyl-β-D-glucosaminide, mean ± SE, n = 3) of AGG\textsubscript{0.8} and AGG\textsubscript{0.2} was 4.00 ± 0.99 nM min\textsuperscript{-1} ml\textsuperscript{-1} and 0.03 ± 0.01 nM min\textsuperscript{-1} ml\textsuperscript{-1}, respectively, with the former being significantly (p < 0.05, Student’s t-test) higher than the latter.

Results of morphological analyses of aggregates are tabulated in Table 1. Neither the long to short axis ratio nor the circularity differed significantly among different size classes for both AGG\textsubscript{0.8} and AGG\textsubscript{0.2} (p > 0.05, ANOVA). The average long to short axis ratios for AGG\textsubscript{0.8} and AGG\textsubscript{0.2} were 0.596

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**Fig. 3.** Relationship between attached bacterial abundance and size (equivalent spherical diameter, ESD) of aggregates AGG\textsubscript{0.8} (■: January; ▼: August; ○: November) and AGG\textsubscript{0.2} (□: January; △: August; ●: November). Values are means ± SE; error bars are those associated with 3 (January) or 4 (August and November) replicated bottles. For each bottle, 10 to 20 aggregates were analyzed. Asterisks indicate the significance of the difference between AGG\textsubscript{0.8} and AGG\textsubscript{0.2} (***p < 0.01, **p < 0.001; Student’s t-test). Solid and dashed lines are linear regressions after double logarithmic transformation for AGG\textsubscript{0.8} and AGG\textsubscript{0.2}, respectively (see ‘Results’).
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0.584, respectively, which did not differ significantly from each other (p > 0.05, Student’s t-test). The average circularities for AGG0.8 and AGG0.2 were 0.072 and 0.044, respectively. The former was significantly larger than the latter (p < 0.001, Student’s t-test).

Fig. 4. Median settling velocity ($W_{50}$ cm s$^{-1}$) of each size class of aggregates AGG0.2 (gray bars) and AGG0.8 (white bars) in (a) January, (b) August, and (c) November experiments. Values are means ± SE of 3 (January) or 4 (August and November) replicated bottles. Asterisks indicate the significance of the difference between AGG0.8 and AGG0.2 (***p < 0.001; Student’s t-test)

Fig. 5. Relationship between porosity and size (equivalent spherical diameter, ESD) of aggregates AGG0.8 (■) and AGG0.2 (□). Data were collected during the November experiment. Values are means ± SE of 4 replicate bottles. Asterisks indicate the significance of the difference between AGG0.8 and AGG0.2 (***p < 0.001; Student’s t-test) for each size class. Fitted curves are quadratic regressions after double logarithmic transformation (see ‘Results’)

DISCUSSION

Use of model aggregates to examine aggregate settling velocity

The model aggregates exhibited the key features of natural marine aggregates: they were spontaneously assembled micro-gels that are abundant in marine waters (Chin et al. 1998, Verduzo 2012); they were stainable by Alcian Blue, a critical feature that characterizes TEP (Allredge et al. 1993); they contained sulfate and amino polysaccharides, which are the major constituents of TEP (Zhou et al. 1998, Pas- sow 2002); and their monomeric constituents, fucose and glucosamine, are ubiquitous and can be abundant in marine waters (Borch & Kirchman 1997, Myklestad et al. 1997). Furthermore, our experiments modeled the fundamental features of microbe-aggregate interactions in marine systems, i.e. colonization, growth, and enzymatic cleavage of polymers (Azam & Malfatti 2007, Nagata 2008). Ectoenzymatic activities cleaving fucoidan and chitin have been detected in seawater (Gooday 1990, Poulicek et al. 1998, Ziervogel & Arnosti 2008), indicating that the model aggregates serve not only as attachment surfaces but also as carbon and nutrient sources for microbes. In fact, in our experiments, AGG0.8 were densely colonized by bacteria, with an abundance of 0.050 to 0.074 cells μm$^{-2}$. This is within the higher range of bacterial abundance reported to be associated with marine TEP within a similar size range of aggregates (0.006 to 0.079 cells μm$^{-2}$; Schuster &
The average cell volume of bacteria associated with AGG0.2 was relatively large (0.46 μm³) and differed little from that of AGG0.8, suggesting that a fraction of small bacteria initially contained in the 0.2 μm filtered seawater became larger during their growth on the substrate-rich aggregates. The bacterial growth on AGG0.8 was accompanied by a substantial increase in ectoenzymatic activity, supporting the notion that bacteria cleaved polymer matrices during colonization and growth. Thus, our model aggregates represent a class of carbohydrate-rich aggregates that harbor dense bacterial communities. The following discussion will focus on the specific mechanisms underlying the regulation of model aggregate settling velocity.

Possible mechanisms of the reduction of aggregate settling velocity via bacterial colonization

The W₅₀ values for AGG0.8 were significantly (1.6- to 4.5-fold) lower than those for AGG0.2. Settling velocities of AGG0.8 would have been affected by the adsorption of submicron-sized TEP that existed in the 0.8 μm filtrate but not in the 0.2 μm filtrate. Although submicron-sized TEP are known to be abundant in natural seawaters, their contributions to total TEP volumes are usually low (<1%), with typical volume concentrations being on the order of 0.1 to 1 ppm in coastal waters (Passow 2002). Although this model does not perfectly describe the sinking of fractal organic aggregates in the sea (Logan & Wilkinson 1990), it is useful for considering the different impacts of size, density, and porosity on aggregate settling velocities (De La Rocha & Passow 2007). Because our comparison of v was made for aggregates belonging to the same size category (i.e. D is the same), the difference in v is ascribed to the difference in either ρ' (mean density difference) or ϕ (morphological effects). For the density effect, there are at least 2 potential mechanisms by which attached bacteria can reduce ρ'. First, bacterial growth may result in the replacement of a certain volume of polysaccharide matrices in aggregates by bacterial cells, which may then result in a reduction in ρ' (bacterial density is generally lower than polysaccharide density). Second, bacteria may increase the ε of aggregates due to the cleavage of polymers.

To examine the mechanisms underlying this reduction in settling velocity, we used the Stokes model (Oliver et al. 1981) as a theoretical framework to relate the aggregate settling velocity (v, cm s⁻¹; hereafter, W₅₀ is referred to as v for the theoretical considerations in this paragraph) to aggregate properties including size (D, the diameter of the particle in cm), mean density (ρ', g cm⁻³), and morphology (ϕ, a dimensionless coefficient of form resistance):

\[
v = \frac{g D^2 (\rho' - \rho_0)}{18 \mu \phi}
\]  

where g is gravitational acceleration (cm s⁻²), μ is the viscosity of seawater (g cm⁻¹ s⁻¹), and ρ₀ is the density of seawater (g cm⁻³). Although this model does not perfectly describe the sinking of fratial organic aggregates in the sea (Logan & Wilkinson 1990), it is useful for considering the different impacts of size, density, and porosity on aggregate sinking velocities (De La Rocha & Passow 2007).

Table 1. Long to short axis ratio and circularity of aggregates AGG0.8 and AGG0.2. The data were collected during the January experiment. Values are means ± SE of triplicate bottles (n = 3) for the different size classes. For each bottle, 40 to 71 particles were analyzed. ESD: equivalent spherical diameter

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Table 2. Equivalent spherical diameter (ESD) of aggregates AGG0.8 and AGG0.2. The data were collected during the January experiment. Values are means ± SE of triplicate bottles (n = 3) for the different size classes. For each bottle, 40 to 71 particles were analyzed. ESD: equivalent spherical diameter
where \( \rho_s \) (g cm\(^{-3}\)) is the density of the solid fraction consisting of polysaccharide matrices with a density of \( \rho_{\text{pol}} \) (g cm\(^{-3}\)) and bacterial cells with a density of \( \rho_{\text{bac}} \) (g cm\(^{-3}\)). \( \varepsilon \) is the fraction of seawater volume relative to the total aggregate volume, and \( \alpha \) is the fraction of bacterial cell volume relative to the total solid volume of the aggregate. Eqs. (2–4) can be rewritten as follows:

\[
v = \frac{g D^2 (1-\varepsilon) [(1-\alpha) \rho_{\text{pol}} + \alpha \rho_{\text{bac}} - \rho_0]}{18 \mu \varphi}
\] (5)

This model assumes that the aggregates are non-permeable (there is no advective flow through the aggregates), which is consistent with the suggestion that marine aggregates have very low permeability (Ploug & Passow 2007, Kindler et al. 2010). Clearly, this assumption does not hold for all aggregate types in marine environments. In fact, for fast-settling, large aggregates (with diameters on the order of a few mm) containing ballast particles, apparent diffusivity within the aggregates has been reported to be high (Ploug et al. 2008). However, for the small (diameters 62 to 119 \( \mu \)m), slowly settling aggregates examined here, it is probably safe to assume that permeability barely influenced aggregate settling velocities (Logan & Hunt 1987).

Using Eq. (5) and the parameters obtained in the present study (\( \epsilon \), \( \alpha \), and \( \rho_{\text{bac}} \)) and from the literature (Table 2), we examined the relative contribution of 2 effects of attached bacteria, i.e. \( \alpha \) and \( \varepsilon \), to the reduction in the settling velocity of aggregates. Because the morphological effect (\( \varphi \)) was unknown, we first estimated the \( \varphi \) value for which the predicted \( v \) becomes equal to the observed \( v \) for each size category of AGG\(_{0.2}\). Then, the average (±1 SE) \( \varphi \) for different size categories (2.80 ± 0.13, \( n = 5 \)) was used as a parameter of the morphological effect for AGG\(_{0.2}\) (parameter set 1). Next, by assuming that \( \varphi \) is constant (= 2.80), we determined to what extent the \( v \) for AGG\(_{0.2}\) changes with increases in \( \alpha \) and \( \varepsilon \) to the levels that we determined for AGG\(_{0.8}\) (parameter sets 2 and 3). The results revealed that the increase in \( \alpha \) (parameter set 2) resulted in only a 4.9% (range: 3.0 to 7.1%) reduction in \( v \) (Table 3), which explained a small fraction of the observed difference in \( v \) between AGG\(_{0.8}\) and AGG\(_{0.2}\) (71.2%). Note that the extent of the reduction in \( v \) with increasing \( \alpha \) might be overestimated because our estimate of bacterial density would be too low (see ‘Results’). In contrast, increases in both \( \alpha \) and \( \varepsilon \) (parameter set 3) resulted in a large reduction of \( v \), with an average reduction of 75.4% (range: 74.4 to 76.4%) (Table 3). These results indicate that the \( \varepsilon \) effect was mainly responsible for the difference in \( v \) between AGG\(_{0.2}\) and AGG\(_{0.8}\). Note that the predicted \( v \) with parameter set 3 was slightly lower (by 0.9 to 6.3% depending on size categories) than the observed \( v \) (Table 3). This might be due to differences in morphology (hence \( \varphi \)) between AGG\(_{0.2}\) and AGG\(_{0.8}\). Consistent with this notion, our results indicated that the circularity of AGG\(_{0.8}\) was larger

### Table 2. Summary of the parameters used for the prediction of aggregate settling velocity (\( v \)) according to the model described by Eq. (5). *Asterisks indicate the choice of parameters for each parameter set

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Units</th>
<th>Parameter set 1</th>
<th>Parameter set 2</th>
<th>Parameter set 3</th>
<th>Parameter set 4</th>
<th>Values</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>( D )</td>
<td>( \mu )m</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>62, 73, 86, 101, 119</td>
<td>Diameter of the aggregate</td>
</tr>
<tr>
<td>( g )</td>
<td>m s(^{-2})</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td></td>
<td>9.8</td>
<td>Gravitational acceleration</td>
</tr>
<tr>
<td>( \mu )</td>
<td>kg m(^{-1}) s(^{-1})</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td></td>
<td>0.00099</td>
<td>Viscosity of seawater at 23°C and salinity 29 (Fofonoff &amp; Millard 1983)</td>
</tr>
<tr>
<td>( \rho_0 )</td>
<td>g cm(^{-3})</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td></td>
<td>1.019</td>
<td>Density of seawater at 23°C and salinity 29 (Fofonoff &amp; Millard 1983)</td>
</tr>
<tr>
<td>( \rho_{\text{pol}} )</td>
<td>g cm(^{-3})</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td></td>
<td>1.6</td>
<td>Density of polysaccharide matrices; Rickwood (1984), Harding (1995)</td>
</tr>
<tr>
<td>( \rho_{\text{bac}} )</td>
<td>g cm(^{-3})</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td></td>
<td>1.064</td>
<td>Density of bacterial cells, determined for bacteria recovered from AGG(_{0.8}) (see ‘Discussion’)</td>
</tr>
<tr>
<td>( \varepsilon ) (AGG(_{0.2}))</td>
<td>–</td>
<td>*</td>
<td>*</td>
<td>–</td>
<td></td>
<td>0.72–0.93</td>
<td>Porosity (( \varepsilon )) for each size class was derived using the empirical relationship between ( D ) and ( \varepsilon ) (see ‘Results’)</td>
</tr>
<tr>
<td>( \varepsilon ) (AGG(_{0.8}))</td>
<td>–</td>
<td>–</td>
<td>*</td>
<td>*</td>
<td></td>
<td>0.93–0.98</td>
<td>Porosity (( \varepsilon )) for each size class was derived using the empirical relationship between attached bacterial abundance and ( D \times ) average bacterial cell volume by total aggregate volume determined by the Coulter counter method (see ‘Results’)</td>
</tr>
<tr>
<td>( \alpha ) (AGG(_{0.2}))</td>
<td>–</td>
<td>*</td>
<td>–</td>
<td>–</td>
<td></td>
<td>0.022–0.004</td>
<td>( \alpha ) for each size class was calculated by dividing total bacterial volume (= bacterial abundance derived using the empirical relationship between attached bacterial abundance and ( D \times ) average bacterial cell volume) by total aggregate volume determined by the Coulter counter method (see ‘Results’)</td>
</tr>
<tr>
<td>( \alpha ) (AGG(_{0.8}))</td>
<td>–</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td></td>
<td>0.036–0.080</td>
<td></td>
</tr>
<tr>
<td>( \varphi ) (AGG(<em>{0.2})) or AGG(</em>{0.8})</td>
<td>–</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td></td>
<td>2.80</td>
<td>Dimensionless coefficient of form resistance</td>
</tr>
<tr>
<td>( \varphi ) (AGG(_{0.8}))</td>
<td>–</td>
<td>–</td>
<td>*</td>
<td></td>
<td></td>
<td>2.42</td>
<td></td>
</tr>
</tbody>
</table>
than that of AGG0.2. The difference between the predicted \( v \) and the observed \( v \) for AGG0.8 was minimized when \( \phi \) was 2.42 (parameter set 4; Table 3), yielding an average reduction value (71.6%) close to the observed value (71.2%).

The derivation of \( \phi \) and the examination of other parameters using the modified Stokes model (Eq. 5) have limitations, especially regarding the simplification of particle geometry. Our model lacks explicit representation of aggregate fractal dimension, which is an important geometric property influencing aggregate settling velocities (Logan & Wilkinson 1990, Vahedi & Gorczyca 2012). Keeping these limitations in mind, we stress that the estimated \( \phi \) would reflect not only the form resistance but also other hydrodynamic forces not examined here (Logan & Wilkinson 1990). Nonetheless, the above results concerning the sensitivity of predicted \( v \) to different parameter settings can be used to infer the principal mechanism by which \( v \) was reduced as a consequence of bacterial colonization. We suggest that the increase in \( \varepsilon \), due to bacterial enzymatic cleavage of aggregate matrices, was mainly responsible for the lower \( v \) for AGG0.8 relative to AGG0.2. Bacterial cell volume (\( \alpha \)) and morphology (\( \phi \)) also contributed to the change in \( v \), although their contributions were not as large as that of porosity.

### Implications for material cycling in marine environments

The aggregate settling velocities that we determined in the present study (0.003 to 0.014 cm s\(^{-1}\)) are within the range of settling velocities reported in the literature for marine particles of similar size (approximately 50 to 100 \( \mu \)m) classes (0.002 to 0.149 cm s\(^{-1}\); Fennessy et al. 1994, Christiansen et al. 2002, Xia et al. 2004). These particles, which are within the smaller size range of marine settling particles (Simon et al. 2002, McDonnell & Buesseler 2012), are abundant in seawater and can account for a large fraction (up to 20%) of total carbon sinking fluxes, as recently estimated for the subtropical Sargasso Sea (McDonnell & Buesseler 2012). In addition, smaller sinking particles, especially sticky microgels, can serve as the precursors of larger aggregates (Verdugo 2012), indicating that changes in settling velocities of smaller particles can alter the depth-dependent patterns of larger aggregate formation in water columns. Considering the timescale of bacterial colonization and growth on fresh aggregates in the upper ocean (day) and the length scale of small aggregates settling at this timescale (10 to 100 m), bacterial porosity enhancement can be an effective mechanism of suppressing the export of organic carbon and other bioelements to depth layers below the euphotic zone. We also note that changes in the porosity of aggregates due to attached bacterial action have implications for the regulation of fluid exchange between interstitial spaces and the ambient water in aggregates. Kindler et al. (2010) suggested that more porous aggregates can retain water for longer at density interfaces because of diffusion-limited fluid entrainment in the interstitial spaces of aggregates. Bacterial enhancement of aggregate porosity can be an important mechanism that promotes aggregate accumulation at density interfaces, a prominent phenomenon widely observed in oceanic environments (Allredge & Gotschalk 1988, MacIntyre et al. 1995).

In addition, bacterial porosity enhancement may result in a higher retention of dissolved organic carbon in the interstitial spaces of aggregates, which may accelerate the downward delivery of dissolved organic carbon mediated by settling aggregates (Allredge 2000).
Natural aggregates in marine waters are complex and variable mixtures of organic and inorganic components. They are often composed of a variety of dense source particles, such as diatom frustules and dust (Ploug et al. 2008, Iversen & Ploug 2010). Depending on the nature and compositions of the aggregates, the extent of bacterial effects on the aggregate settling velocities might deviate from the present results. For example, bacterial effects on the settling velocities would be less pronounced for the aggregates with a low content of polysaccharides and a high content of mineral particles because bacterial enzymatic cleavage of polymers may hardly modify bulk physical properties of such aggregates. It is important for future studies to examine the relationship between aggregate compositions and the extent and mechanisms of bacterial effects on aggregate settling velocities.

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